A96

BA DI6

Europäisches Potentomt
Europeun Patent Giffee
Ciffee europeen des brovets

11 Publication number:

0 039 **245** 2

(B)

## EUROPEAN PATENT APPLICATION

Application .number: 63301470.7

(1) Int. Cl.2: A 61 K 45/02

Date of filing: 16.03.03

STABLE INTERFERON BETA CATUPSA. + WITH HIGHLY PURIFIED INTERFERON BETA IN BUFFER CONTG. POLY VINYL-PYRROLIDONE

- (3) Priority: 17.03.62 IL 65277
- ② Cote of publication of application: 21.03.03 DuCartin 02/03
- Designated Commenting States:
  AT DE CI DE FR GD IT U NL SE

- Applicant: DVTER-YEDA LTD.
   Kiryat Welzmann
   Ness-Zionejill
- (2) Inventor: Symbolista, Samuel 3 Hazionut Circon Jerusalem(IL)
- (14) Representative: Sewors, Lowrence Peter et al, PAGE, White & FARRER 5 Plough Place New Fatter Lane
  London EC4A 1HY(G3)

1733

772523

Craydon Printing Company Ltd

(7

A stable interferon beta composition and a method of stabilizing interferon beta.

<sup>)</sup> This invention relates to a method of stabilizing Interference  $\beta$  (Human Fibroblast Interferen (HFIF)), wherein a highly purified Imerferon  $\beta$  solution, admixed with known exciplents therefor, is dialyzed against an exercise buffer at pH 3.5 for about 43 hours. The resulting Interferon  $\beta$  solution is admixed with from 0.5% to 10% vervalume of polyvinyl symplectic prior to or following filtration through a sterile filter, dispensed into glass vials, lyophilized, and the vials are scaled in vecus and stored at 4°C.

by means of vinyl pyrrolidone polymer, hereinafter designated P.V.P., a polymer which has been known for a long time exclusively as a clarifying agent in wines and as a dispersing and suspending agent for pharmaceutical compositions. P.V.P. has molecular weights ranging from 10,000 to 700,000 and it is marketed under trademarks such as POVIDONE et al, (see "The Merck Index", 9th edition, page 7485 under No. 7498).

This invention relates to a novel stable Interferon 3 composition comprising a buffered solution of highly purified Interferon 3 and conventional excipients, said solution being stabilized by 0.5 to 10% wt/volume of polyvinyl pyrrolidone.

This invention also relates to a method of stabilizing
15 Interferon β, wherein a highly purified Interferon β
solution, admixed with known excipients therefor, is
dialysed against an acetate buffer solution, the
resulting Interferonβ solution is admixed with from 0.5%
to 10% wt/volume of polyvinyl pyrrolidone prior to or
20 following filtration through a sterile filter, dispensed
into glass vials, lyophilized, and the vials are sealed
in vacuo and stored at 4°C. The dialysis is preferably
continued for about 48 hours.

The preferred excipients are mannitol and human serum albumin (MSA). The acetate buffer used contains sodium acetate and sufficient acetic acid to adjust the pH to 3.5. P.V.P. marketed as POVIDONE, having a molecular weight of about 50,000, is the preferred stabilizer, but P.V.P., having lower or higher molecular weights has also proved to be highly effective as a stabilizer.

The preparation of the preferred inventive Interferon  $\beta$  composition will now be described in the following example. The preparation of the preferred inventive Interferon  $\beta$ 

10 lts of aqueous acctate buffer solution having a pH=3.5 are prepared by dissolving 21.6 cc of acetic acid and 4.02 gms of sodium acetate in the required volume of distilled water.

The inner surface of a sterile dialysis bag is wetted with sufficient concentrated human serum albumin to 10 result in a 1% concentration in a highly purified Interferon β solution, having a specific activity of about 10<sup>7</sup> international units per mg of protein, which is subjected to dialysis therein.

The resulting solution is dialysed against the acetate 15 buffer of pH 3.5 and at a temperature of  $4^{\circ}\text{C}$  for about 48 hours at a ratio of 1:100 Interferon \$\beta\$ solution to buffer solution with a change of the buffer solution after 24 hours.

The dialysed Interferon \$\beta\$ preparation is admixed with 20 mannitol 0.5 wt/volume final concentration and with P.V.P. at a 2% final concentration approximately prior to or following filtration through a sterile filter, previously imprognated with sufficient concentrated human serum albumin to raise the albumin concentration in the 25 filtrate to 2% wt/volume. The filtrate is collected in a sterile bottle.

The P.V.P. concentration is then finally adjusted to 2% wt/volume and the concentration of mannitol to 0.5% wt/volume, if necessary. The final volume of the solution 30 is adjusted with sterile acetate buffer.

1730

2 cc each of the solution obtained are dispensed into sterile glass vials by means of a sterile Cornwall syringe, followed by lyophilization and the vials are then scaled in vacuo and stored at 4°C. The contents of the vials are resuspended by the addition of 2 cc of bidistilled sterile water.

The composition of the final product per vial is as follows:

	Sodium Acetate AG	0.4	ang m	
10	Sodium Chloride AC	1.3	ngn	
	Human Serum Albumin Fraction	V 40.0	மஜம	
	Hannitol AG	10.0	egt.	
	PVP - Stabilizer	40.0	±∑	
	Muman Fibroblast Interferon	1.0 x	: 10 <sup>6</sup> I.U.	(approximately)

- The effectiveness of P.V.P. of different molecular weights and in different concentrations on the stability of Interferon  $\beta$  in its compositions will now be illustrated by the following Tables 1 to 6 of which: Table (1) illustrates the effect of P.V.P. of molecular weight 24,000 at concentrations ranging from 0.5% to 5% on the
- 24,000 at concentrations ranging from 0.5% to 5% on the stability of Interferon β in its inventive compositions immediately before and after lyophilization, and after storage for 1 to 4 months in sealed vials at 37°C. The data in Tables (2) and (3) illustrate the stability of the
- compositions under identical conditions, using P.V.P.
  of molecular weights 50,000 and 160,000 respectively.
  Comparative data are reported in these tables for
  Interferon \$ compositions without P.V.P. and compositions
  containing sucrose or human serum albumin in various con-
- 30 centrations instead of P.V.P. Tables (4), (5) and (6) illustrate the relationship between the titre of inventive Interferon β compositions and their content of P.V.P.

1707

of the same molecular weights as in Tables (1), (2) and (3):

- (a) immediately after resuspension as hereinbefore described, and,
- 5 (b) after storage of the resuspended compositions for 1 month at 4°C.

Comparative data for Interferon  $\beta$  compositions admixed with sucrose or human serum albumin are again given.

The following data and remarks are essential for the understanding of these tables:

- (a) The Human Fibroblast Interferon used in the compositions was initially purified to a specific activity ranging from 10<sup>6</sup> to 10<sup>7</sup> international units per mg of protein.
- 15 (b) The data in the Tables relating to the titres of Interferon β in admixture with P.V.P. in different concentrations are the averages of 6 titration results and the data are expressed in megaunits per vial.
- (c) The difference in the initial titres of Inter feron β are due to the use of Interferon β from different
   batches which differ somewhat in their specific activity.

It is evident from the data reported in the tables that P.V.P. of different molecular weights have maximal stabilizing effectiveness when used in concentration of from

25 2 to 4% although the use of P.V.P. in concentrations of up to 10% also leads to positive stabilization results. Positive stabilization is also attained using P.V.P. having molecular weights from 10,000 to 700,000.

Othermodifications of the method described hereinbefore 30 are known to the man versed in the art and these are included therein provided that they fall within the ambit of the invention defined in the subsequent claims.

Conc. of PVP	Titre before Lyophilization	Titre after Lyophilization	Titre after 1 Month at 37°C	Titre after 2 Months at 37°C	Titre after 3 Months at 37 C	Titre after 4 Konths at 37 C
Û	j.4	0.8	0.00	0.02	<b>4</b> 0.01	< 0.01
0.5	1.3	1.0	0.8	0.6.	0.2	0.20
::	1.5	1.3	1.2	1.2	1.1	0.0
2.5	1.3	1.3	1.2	1.2	1.1	0.0
3.	1.3		1.1	1.2	1.1	1.1
, o T	1.6	1.4	1.2	1.1	1.0	1.0
· •	1.4	1.0	6:0	1.0	0.8	8.0
0 + Sucrose 5%	1.5	7.0	0.05	0.01	< 0.01	10.0 >
O + Sucrose 103	1.3	8.0	0.04	0.01	< 0.01	70.0
0 + 115A 3\$	1.4	8.0	6.1	0.03	< 0.01	. < 0.01
0 + 11SA 45	1.3	0.0	0.1	0.05	0.02	10.0

772523

Polyvinylpyrrolidone 24000

Titrg basore Titre after Tiure after Tit.3 after Lyophilization Lyophilization 1 Month 2 Houths 2 Houths at 37°C at 37				Table 2			
1.1       0.7       0.06       0.01         1.2       0.9       0.6       0.1         1.1       1.1       1.1       1.1         1.2       1.1       1.2       1.2         1.1       1.2       1.2       1.2         1.1       1.2       1.2       1.0         1.3       1.2       1.2       1.0         1.0       0.9       4.0       0.8       0.02         2.103       1.2       0.8       0.03       0.03       0.03         2.104       1.2       0.8       0.003       0.03       0.03	Conc. of PVP	Titra before Lyophillzation	Titre after Lyophilization	Tiure after 1 Nonth at 37°C	Tites after 2 Houths at 37°C	Titre after 3 Hoaths 2* 370C	Titre after 4 Eouths et 37°C
1.2       0.9       0.6       0.1         1.1       1.1       1.1         1.2       1.1       1.2         1.1       1.2       1.2         1.1       1.2       1.2         1.3       1.2       1.0         1.0       0.9       4.0       0.8         c 55       1.1       0.6       0.05       0.02         c 103       0.08       0.09       0.03       0.03         c 103       0.03       0.03       0.03       0.03	. 0	1.1	0.7	90.0	0.01	< 0.01	< 0.01
1.1 1.1 1.1 1.2 1.2 1.1 1.2 1.2 1.1 1.2 1.2 1.1 1.2 1.2 1.2 1.0 1.0 0.9 4.0 0.8 0.05 0.05 C.01 2.107 1.2 C.01	0.5%	1.2	6.0	9.0	0.1	0.07	0.03
1.2 1.2 1.2 1.2 1.2 1.2 1.1 1.2 1.2 1.2		1.1	1.1	1.1	1.1	1.0	0.8
1.1 1.2 1.2 1.2 1.2 1.0 1.3 1.2 1.0 1.0 1.0 1.0 0.9 1.0 0.8 0.05 0.05 0.05 0.05 0.05 0.05 0.0	2.6	. 1.2	1.1	1.2	1.2	1.1	1.0
1.3 1.2 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0	31	1.1	1.2	1.3	1.2	1.1	6.0
1.0 0.9 4.0 0.8 0.02 < 0 0.1 0.6 0.07 0.02 < 0 0.1 0.1 0.8 0.05 0.05 0.01 < 0 0.01 0.01	4%	1.3	1.2	1.2	1.0	1.2	1.0
2 101 1.1 0.6 0.07 0.02 < C 1.1 1.2 0.8 0.09 0.03 < C 1.1 1.2 0.8 0.09 0.03 < C 1.1 1.2 0.8 0.09 0.03	\$\$	1.0	6.0	€.0	8.0	0.3	0.9
2 101 1.2 0.8 0.05 C.01 < 1.2 1.2 0.8 0.09 0.03 < 1.2	0 + Sucrose St	1.1	9.0	0.07	0.02	< 0.01	< 0.01
1.2 0.8 0.09 0.03	0 + Sucross 101	1.2	8.0	0.05	0.01	< 0.01 ·	< 0.01
	0 + 11SA 3%	1.2	0.8	0.09	0.03	< 6.01	4 0.01
1:1 0.8 0.10 0.03	0 + HSA 4%	1.1	0.8	0.10	0.03	0.01	7 0.01

Polyvinylpyrrolidons 50000

	Q(	30	2	4	?
--	----	----	---	---	---

			2001	>		
Conc of PVP	Titre before Iyophilization	Titre before Titre after Iyophilization Iyophilization	Titre after 1 Month at 37°C	Titre after 2 Nonths at 37 G	Titre after 3 Honths at 37 G	Tite after 4 Fonths at 37 C
20	1.5	. w.o	0.05	0.01	10 1 7	
0.5%	1.4	1.0	50.0	90.0	16.5	10.0 >
**	1.6	1.3	1.0	9.0	Ç. 4	0.03
21	1.4	1.6	1.5		÷ -	٠.0
٠.	1.5	1.5	1.3	1.4		c
4.	1.6	1.4	1.3	1.4		7.7
54	1.7	1.3	1.4.	F. T.	1.3	1.2
0 + Sucrose 5%	5 5 1.7	6.0.	0.04	0.01		1.2
0 + Sucrose 10% 1.3	101 1.3	0.8	0.05	< 0.01	7 0.61	<0.01 0.01
0 + 11SA 38	1.4	0.8	0.03	0.04	. 0.01	Z 0.01
0 + 11SA 4%	1.5	1.0	0.1	0.05	. 0.01	< 0.01 < 0.01

æ	
ပ	
5	
۳	

Conc. of Pvp	Titre of IP With resuspension	Titre of 16 1 Nonth after
		resuspension stored at 4°C
	0.8	< 0.01
• • • • • • • • • • • • • • • • • • •	1.0	0.3
	1.3	9.0
	. 1.3	1.2
	. 1.2	1.1
	1.4	1.2
0 + Sucrose 5th	1.0	1.1
0 + Sucrose 10\$	0.7	< 0.01
0 + 11SA 3%	0.8	< 0.01
0 + 115A 41	. 8.0	< 0.01
	6.0	40.01

Polyvinyl pyrrolidone 24030

•	•	•
	ς	د
٠	_	4
	-	>
	C	:
ı	_	٠

Conc of PVI	Titre of IF with resuspension	Tite of IF 1 Month after resuspension stand
. %0		10 C 10 C 201 2101 2101 2101 2101 2101 2101
,	7.0	< 0.01
	6,0	0.5
5 6 5 6 5 6 5 6 5 6 5 6 5 6 5 6 5 6 5 6	1.1	0.3
72		1.0
5.2	1.2	1.1
ار این از ای این از این ا	7.7	1.2
0 + Siterosa 5%	. v	1.0
0 + Sucrose 10%		< 0.01
0 + 115A 34		< 0.01
0 + 115A 4%	. 0.8	<ul><li>10.0 &gt;</li><li>20.0 i</li></ul>
		. 50.0

Polyvinyl pyrrolidone 50000

Titra of IF 1 Month  ***Ifficial Control of IF 1 Month  ***Ifficial of IF 1 Month  **Ifficial of IF 1 Month  ***Ifficial of IF 1 Month  **Ifficial of IF 1 Month  ***Ifficial of IF 1 Month  **Ifficial of IF 1 Month  ***Ifficial of IF 1 Month  **Ifficial of IF 1 Month  ***Ifficial of IF 1 Mon	Titre of IF  Titre of IF  With resuspension  0.8  1.0  1.0  1.5  1.4  1.5  1.4  1.5  0.9  2.0.01  0.8  1.4  1.5  1.4  1.5  1.7  1.0  0.8  1.0  0.8  1.0  0.9  2.0.01  2.0.01  1.0	c of IF 1 resuspension 0.8 1.0 1.5 1.5 1.4 1.5 0.9 0.9	pa	-:1-
IF	Table 6  Titre of IF with resuspension  0.8  1.0  1.3  1.6  1.5  1.7  0.9  0.8  0.8	Tabi	Titra of IF 1 Month after resuspension stor	2 0.01 C 3 n.8 1.4 1.3 1.4 1.3 2 0.01 2 0.01
<b>3</b>	Table 6  Titre of with res  1.0  1.3  1.5  1.4   0.9   1.0	Tabi	IF uspension	

## CLAIMS:

5

- 1. A stable Interferon β composition comprising a buffered solution of highly purified Interferon β and conventional excipients, said solution being stabilized by 0.5 to 10% wt/volume of polyvinyl pyrrolidone.
  - 2. A composition as claimed in Claim 1, wherein the excipients are mannitol and human serum albumin.
- 3. A composition as claimed in Claim 1 or 2,10 wherein the buffer is an acetate buffer having a ph of 3.5.
  - 4. A composition as claimed in any one of the preceding claims packaged in a glass vial, lyophilized and sealed in vacuo.
- 15 5. A method of stabilizing Interferon  $\beta$ , wherein a highly purified Interferon  $\beta$  solution, admixed with known excipients therefor, is dialysed against an acetate buffer solution, the resulting Interferon  $\beta$  solution is admixed with from 0.5 to 10% wt/volume
- of polyvinyl pyrrolidone prior to or following filtration through a sterile filter, dispensed into glass vials, lyophilized, and the vials are sealed in vacuo and stored at 4°C.
- 6. A stabilized Interferon  $\beta$  composition whenever obtained by the method claimed in Claim 5.